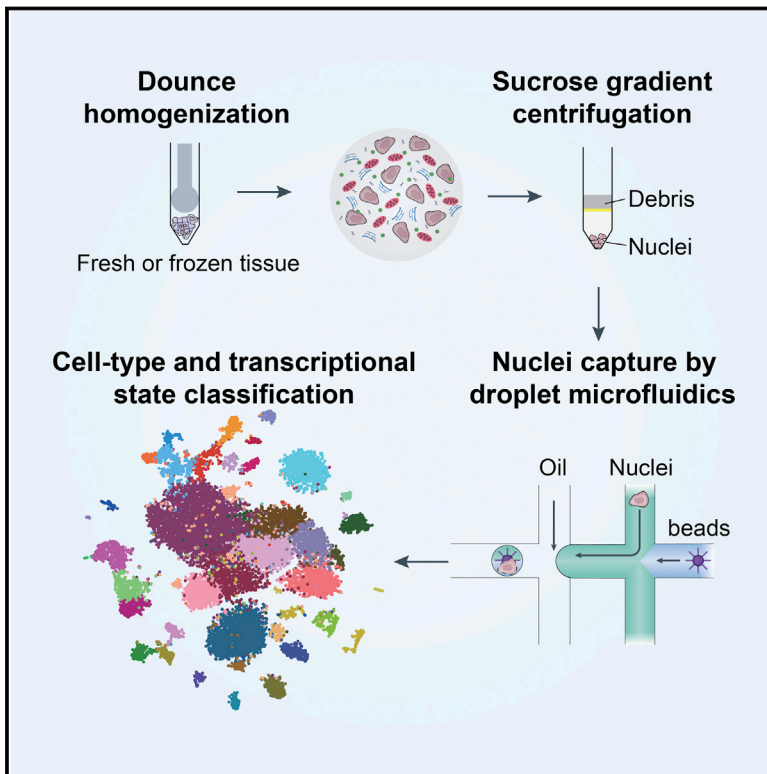


Molecular Cell

Dissecting Cell-Type Composition and Activity-Dependent Transcriptional State in Mammalian Brains by Massively Parallel Single-Nucleus RNA-Seq

Graphical Abstract



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In Brief

Hu et al. developed sNucDrop-seq, a highly scalable single-nucleus RNA-seq method that resolves cell-type heterogeneity and enables the dynamics of transient transcriptional states to be analyzed with single-cell resolution in freshly isolated or frozen mammalian tissues. This allows mapping of seizure-induced transcriptional responses at single-cell resolution in mammalian brains.

Highlights

- We report “sNucDrop-seq,” a massively parallel single-nucleus RNA-seq method
- sNucDrop-seq reveals neuronal subtype composition in adult mammalian brain regions
- sNucDrop-seq enables the dissection of transient transcriptional states
- sNucDrop-seq enables mapping of cell-type-specific response to PTZ-induced seizures



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Dissecting Cell-Type Composition and Activity-Dependent Transcriptional State in Mammalian Brains by Massively Parallel Single-Nucleus RNA-Seq

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SUMMARY

Massively parallel single-cell RNA sequencing can precisely resolve cellular diversity in a high-throughput manner at low cost, but unbiased isolation of intact single cells from complex tissues such as adult mammalian brains is challenging. Here, we integrate sucrose-gradient-assisted purification of nuclei with droplet microfluidics to develop a highly scalable single-nucleus RNA-seq approach (sNucDrop-seq), which is free of enzymatic dissociation and nucleus sorting. By profiling ~18,000 nuclei isolated from cortical tissues of adult mice, we demonstrate that sNucDrop-seq not only accurately reveals neuronal and non-neuronal subtype composition with high sensitivity but also enables in-depth analysis of transient transcriptional states driven by neuronal activity, at single-cell resolution, *in vivo*.

INTRODUCTION

A fundamental challenge in deciphering cell-type composition and cells' functional states in complex mammalian tissues manifests in the extraordinary diversity of cell morphology, size, and local microenvironment. While existing single-cell RNA-seq approaches have proven to be powerful tools for interrogating cell types, dynamic states, and functional processes *in vivo* (Tanay and Regev, 2017), these methods require the preparation of intact, single-cell suspensions from freshly isolated tissues, which is only practical for easily-dissociated embryonic and young postnatal tissues. This requirement poses an even greater challenge for cells with complex morphology, such as mature neurons. Enzymatic treatment not only favors recovery of easily dissociated cell types, but also introduces aberrant transcriptional changes during the whole-cell dissociation process (Lacar et al., 2016; Wu et al., 2017). In addition, skeletal and cardiac muscle cells are frequently multinucleated and are large in size. For instance, each adult mouse skeletal muscle cell contains hundreds of nuclei and is ~5,000 μm in length

and 10–50 μm in width (White et al., 2010). Thus, existing high-throughput single-cell capture and library preparation methods—including isolation of cells by fluorescence-activated cell sorting (FACS) into multi-well plates, sub-nanoliter wells, or droplet microfluidic encapsulation—are not optimized to accommodate these unusually large cells. Isolating individual nuclei for transcriptome analysis is a promising strategy, as single-nucleus RNA-seq methods avoid strong biases against cells of complex morphology and large size (Habib et al., 2016; Lacar et al., 2016; Lake et al., 2016; Zeng et al., 2016) and can be potentially standardized to accommodate the study of various tissues. However, current single-nucleus RNA-seq methods primarily rely on fluorescence-activated nuclei sorting (FANS) (Habib et al., 2016; Lake et al., 2016) or the Fluidigm C1 microfluidics platform (Zeng et al., 2016) to capture nuclei; thus, they cannot easily be scaled up to generate a comprehensive atlas of cell types in a given tissue, much less a whole organism.

DESIGN

An ideal solution to increase the throughput of single-nucleus RNA-seq is to integrate nucleus purification with massively parallel single-cell RNA-seq methods such as Drop-seq (Macosko et al., 2015), InDrop (Klein et al., 2015), or commercial platforms such as 10 \times Genomics (Zheng et al., 2017). However, single-nucleus RNA-seq is currently not supported on these droplet microfluidics platforms. Inefficient lysis of nuclear membranes and/or cellular debris contamination might contribute to this failure. Historically, nuclei of high purity can be isolated from solid tissues or from cell lines with fragile nuclei by centrifugation through a dense sucrose cushion to protect nucleus integrity and strip away cytoplasmic contaminants. The sucrose-gradient ultracentrifugation approach has been adapted to isolate neuronal nuclei for profiling histone modifications, nuclear RNA, and DNA methylation at genome scale (Johnson et al., 2017; Lister et al., 2013; Mo et al., 2015). Here, we develop “sucrose-gradient-assisted single-nucleus Drop-seq” (sNucDrop-seq), a method that enables highly scalable profiling of nuclear transcriptomes at single cell resolution by integrating sucrose-gradient ultracentrifugation-based nucleus purification with droplet microfluidics.

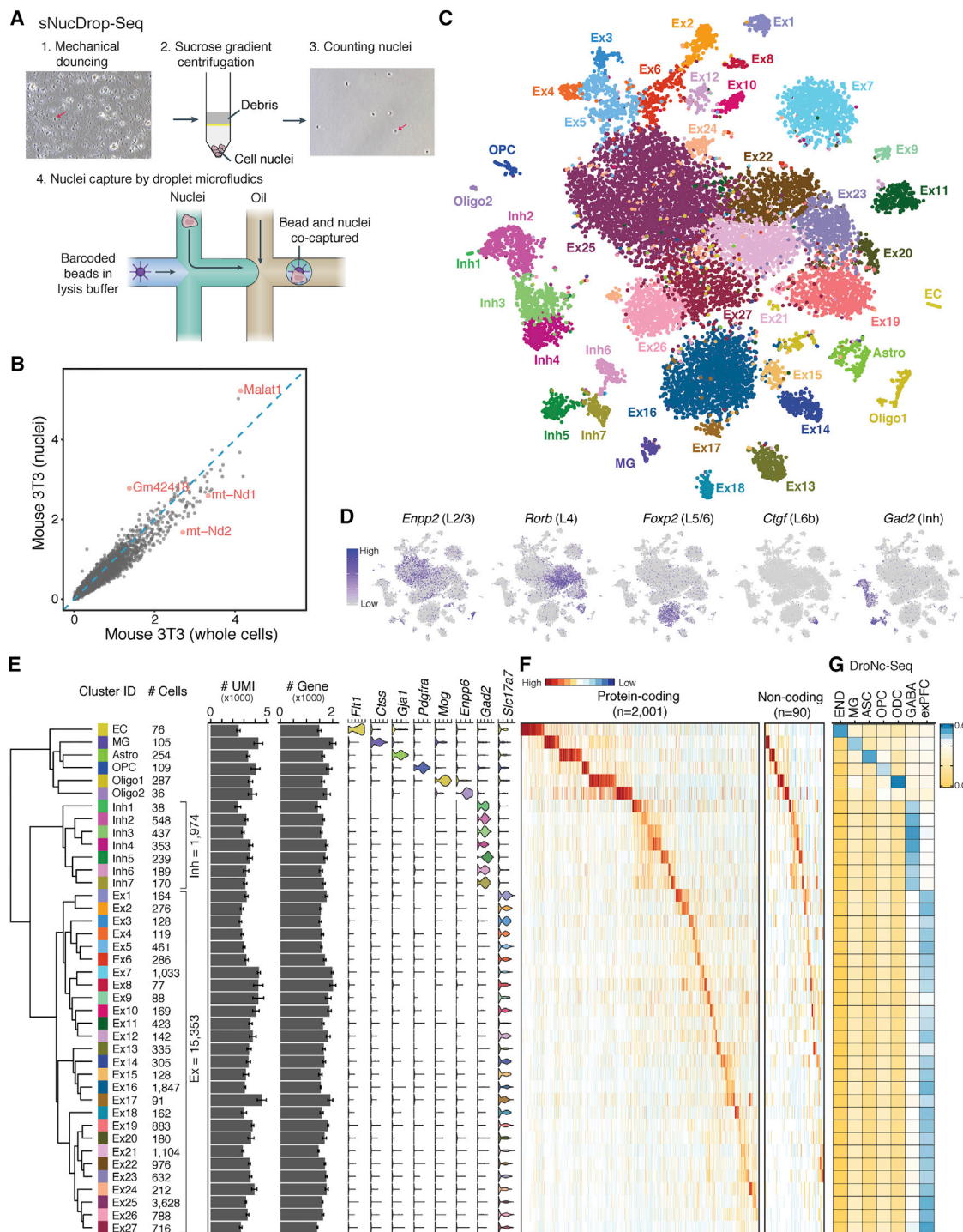


Figure 1. sNucDrop-Seq: A Massively Parallel Single-Nucleus RNA-Seq Method

(A) Overview of sNucDrop-seq. Red arrows indicate representative nuclei before or after sucrose-gradient centrifugation.

(B) Scatterplot comparing the average expression levels detected in NIH3T3 nuclei (y axis, by sNucDrop-seq) and cells (x axis, by Drop-seq). Red dots mark representative genes preferentially enriched in either nuclei or whole cells.

(C) Visualization by tSNE plot of clustering of 18,194 single-nucleus expression profiles from adult mouse cortices ($n = 17$ mice). Ex, excitatory neurons; Inh, inhibitory neurons; Astro, astrocytes; OPC, oligodendrocyte precursor cells; Oligo, oligodendrocytes; MG, microglia; EC, endothelial cells.

(D) Marker gene expression, shown by recoloring nuclei according to the expression level and projecting onto the tSNE plot in Figure 1C.

(E) Dendrogram showing relatedness of cell clusters, followed by (from left to right) cluster identification (ID), cell number per major cell type, UMIs per cluster (mean \pm SEM), number of genes detected per cluster (mean \pm SEM), and violin plots showing select marker genes for major cell types.

(legend continued on next page)

RESULTS

Validation of sNucDrop-Seq

To test whether this nucleus purification method supports single-nucleus RNA-seq analysis, we isolated nuclei from cultured cells as well as freshly isolated or frozen adult mouse brain tissues through dounce homogenization followed by sucrose-gradient ultracentrifugation (Figure 1A and Figure S1A). After quality assessment and counting of nuclei, we performed emulsion droplet barcoding of the nuclei and library preparation. We found that the Drop-seq platform yielded high-quality cDNA libraries from both whole cells and nuclei (Figure S1B).

We next validated the specificity of sNucDrop-seq with species-mixing experiments using nuclei isolated from cultured mouse and human cells. This analysis indicates that the rate of coencapsulation of multiple nuclei per droplet (~2.6%) is comparable to standard Drop-seq (Figure S1C). To assess the sensitivity of sNucDrop-seq, we performed shallow sequencing of cultured mouse 3T3 cells at either single-cell (with Drop-seq: detecting on average 3,325 genes with ~25,000 reads per cell for 1,160 cells with >800 genes detected) or single-nucleus (with sNucDrop-seq: detecting on average 2,665 genes with ~23,000 reads per nucleus for 1,984 nuclei with >800 genes detected) resolution (Figure S1D). With standard Drop-seq microfluidics devices and flow parameters, the capture rate of sNucDrop-seq (1.9%, 1,829/95,000 barcoded beads) is comparable to that of Drop-seq (1.5%, 1,160/77,000 barcoded beads). Comparative analysis reveals that mitochondria-derived RNAs (e.g., *mt-Nd1*, *mt-Nd2*) and nucleus-enriched long-noncoding RNAs (e.g., *Malat1*) were enriched in reads derived from Drop-seq and sNucDrop-seq, respectively (Figure 1B). Thus, integrating sucrose-gradient centrifugation-based nuclear purification with the Drop-seq microfluidic platform and workflow may support massively parallel single-nucleus RNA-seq.

Application of sNucDrop-Seq to Adult Mouse Cortex

To demonstrate the utility of sNucDrop-seq in studying complex adult tissues, we analyzed nuclei isolated from adult mouse cerebral cortex (Table S1). The average expression profiles of single nuclei from two biologically independent replicates were well correlated ($r = 0.993$; Figures S1E and S1F). Out of reads uniquely mapped to the genome (78% of all reads), 76% of reads were aligned to the expected strand of genic regions (25% exonic and 51% intronic), and the remaining 24% to intergenic regions or to the opposite strand of annotated genic regions. The relatively high proportion of intronic reads is similar to a previous single-nucleus RNA-seq study of human cortex (49%) (Lake et al., 2016), reflecting the enrichment of nascent, preprocessed transcripts in nuclei. Because most exonic (91%) and intronic (86%) reads were mapped to the expected strand of annotated transcripts, we retained both exonic and intronic reads for downstream analyses. After initial quality filtering

(>800 genes detected per nucleus), we retained 20,858 nuclei (15,471 uniquely mapped reads per nucleus), detecting, on average, 3,464 transcripts (unique molecular identifiers [UMIs]), and 1,662 genes per nucleus. After correcting for batch effects, we identified highly variable genes, and determined significant principal components (PC). We then performed graph-based clustering and visualized distinct groups of nuclei using non-linear dimensionality reduction with spectral *t*-distributed stochastic neighbor embedding (tSNE). After removing non-cortical cells (~7%, mostly striatal inhibitory neurons) and data points contributing toward potential noise (see STAR Methods), our analysis segregated 18,194 cortical nuclei into 40 distinct clusters (Figure 1C). Each cluster contains nuclei from multiple animals, indicating the transcriptional identities of these cell-type-specific clusters are reproducible across biological replicates (Figure S2A).

On the basis of known markers for major cortical cell types, we identified 27 excitatory neuronal clusters (Ex 1–27: *Slc17a7*⁺), 7 inhibitory neuronal clusters (Inh 1–7: *Gad2*⁺), and 6 non-neuronal clusters (astrocytes [Astro: *Gja1*⁺], oligodendrocyte precursor cells [OPC: *Pdgfra*⁺], oligodendrocytes [Oligo1: *Mog*⁺; Oligo2: *Enpp6*⁺], microglia [MG: *Ctss*⁺], and endothelial cells [EC: *Flt1*⁺]) (Figures 1C–1F). Consistent with previous studies (Lake et al., 2016; Madisen et al., 2015; Zeisel et al., 2015), known layer-specific marker genes (Layer (L)2 and L3: *Enpp2*; L4: *Rorb*; L6a: *Foxp2*; L6b, *Ctgf*) can be readily detected in specific excitatory neuronal clusters (Figure 1D), revealing the anatomic locations of these glutamatergic excitatory neuronal subtypes. We also uncovered all major subclasses of cortical inhibitory neurons named for the neurochemical markers they express: somatostatin-expressing (*Sst*⁺: cluster Inh1 and Inh2) cells, parvalbumin-expressing (*Pvalb*⁺: cluster Inh3 and Inh4) cells, vasoactive intestinal peptide-expressing (*Vip*⁺: cluster Inh6) cells, and cells that express 5-hydroxytryptamine receptor 3A (*Htr3a*) but lack *Vip* expression (*Tnfrsf813*⁺/*Sema3c*⁺/*Vip*[−]: cluster Inh5 and Inh7) (Figures 1C–1E and S2B). This unbiased sampling strategy captured enough cells to resolve heterogeneity among non-neuronal cell types present in relatively low abundance in the adult cortex (Figures 1E and S2B), including two oligodendrocyte subtypes (Oligo1: *Mog*⁺/*Enpp6*[−]; Oligo2: *Mog*[−]/*Enpp6*⁺) recently identified through single-cell deep sequencing of full-length mRNAs (Tasic et al., 2016; Figure S3C). Finally, the cell types and their signatures from sNucDrop-seq were comparable to those obtained with DroNc-seq (a recently published approach similar to sNucDrop-seq) of mouse prefrontal cortex (Habib et al., 2017; Figure 1G).

In addition to subtype-specific protein-coding marker genes (Figure 1F), we have identified a list of long noncoding RNAs that are specifically expressed in distinct cell clusters (Figures 1F and S2B). For instance, *1700016P03Rik* is specifically detected in clusters Ex17 and Ex24; this acts, mainly, as a primary, noncoding transcript encoding two microRNAs (*Mir212* and

(F) Heatmap showing expression of cell-type-specific protein-coding and long noncoding RNA markers in clusters defined in Figure 1E.

(G) Cell-type-specific expression signatures (identified by sNucDrop-seq) agree with previously published work. Pairwise correlations of the average expression for the genes in each cell-type signature defined by sNucDrop-seq and cell types defined by DroNc-seq in the mouse prefrontal cortex (Habib et al., 2017). DroNc-seq clusters: exPFC, excitatory neurons; GABA, inhibitory neurons; ASC, astrocytes; OPC, oligodendrocyte precursor cells; ODC, oligodendrocytes; MG, microglia; END, endothelial cells.

Mir132), which are regulated by neuronal activity (Aten et al., 2016; Nudelman et al., 2010), raising the possibility that Ex17 and Ex24 clusters are specifically associated with activity-induced transcriptional states. Cell-type-specific noncoding RNA marker genes have also been identified for inhibitory neuronal subtypes (e.g., *Dlx1as* for Inh7) and non-neuronal cells (e.g., *4933406/18Rik* for MG). The identification of both protein-coding and noncoding transcripts as cell-type-specific markers highlights the potential of sNucDrop-seq in exploring the emerging role of noncoding RNAs at single-cell resolution *in vivo*.

We next explored whether aggregation of our single-nucleus RNA-seq data into subtype-specific transcriptome profiles, as a proof of concept, might enable analysis of differential mRNA processing in a cell-type-specific manner. Using a probabilistic model that quantitates the expression level of alternatively spliced genes (MISO) (Katz et al., 2010), we identified 263 differential exon processing events through pairwise comparison of cell-type-specific transcriptome profiles. In agreement with a previous single-cell RNA-seq study (Tasic et al., 2016), Syntaxin binding protein 1 (*Stxbp1*) mRNA exhibited differential processing of a specific exon amongst excitatory and inhibitory neurons (Figure S2C). In particular, additional heterogeneity in the usage of this exon was detected among major inhibitory neuronal subtypes (*Sst*⁺ versus *Pvalb*⁺ in Figure S2C). Our cell-type-specific analysis also revealed differentially processed exon usage between abundant cortical cell-types such as excitatory neurons and relatively rare non-neuronal cells (e.g., *Macf1* in Figure S2C). Thus, despite the shallow sequencing depth and 3' bias, sNucDrop-seq allows a population of highly heterogeneous cells to be analyzed together to discover cell-type-specific signatures of mRNA processing.

sNucDrop-Seq Reveals Composition of Cortical Inhibitory Neurons

GABAergic interneurons are highly diverse in terms of morphology, connectivity, and physiological properties (Kepecs and Fishell, 2014), but the relative composition of these neurons, particularly those of low abundance in the cortex, is not well established. To accurately measure the subtype composition and validate the specificity of cortical inhibitory neuronal subtypes identified by sNucDrop-seq, we performed subclustering on cortical inhibitory neuronal nuclei (Inh1–Inh7 in Figure 2A) together with noncortical nuclei isolated from dorsal striatum (>95% striatal cells are GABAergic neurons), identifying 17 subclusters (Figure 2A). Based on expression patterns of known marker genes, we first segregated these subclusters into cortical interneuron (clusters A–I in Figure 4A: *Gad1*⁺/*Gad2*⁺/*Meis2*[−]) and noncortical (clusters in grey: *Meis2*⁺) clusters. Consistent with their striatal origin, many noncortical cells express *Ppp1r1b* (also known as *DARPP-32*) (Figures 2A and 2B), a marker gene indicative of medium spiny neurons (MSNs, D1-type) in the striatum. Because sNucDrop-seq samples nuclei in proportion to cells' abundance in their native environment, this approach enables direct measurement of subtype composition. This analysis identified *Pvalb*-expressing subtypes (clusters D, E, and F: 40.2%) and *Sst*-expressing subtypes (clusters G, H, and I: 31.4%) as two major groups of cortical interneurons (Figures 2C and 2D), in complete accordance with previous observations

derived from *in situ* hybridization (ISH)- or immunostaining-based analysis of mouse neocortex (Rudy et al., 2011). Beyond major interneuron subtypes, we identified 10.8% of cortical interneurons as an *Ndnf*-expressing subtype (cluster A), 8.3% as a *Vip*-expressing subtype (cluster B), and 9.3% as a synuclein gamma (*Sncg*)-expressing subtype (cluster C) (Figures 2C and 2D). On the basis of combinatorial expression of known marker genes, interneuron subtypes identified by sNucDrop-seq parallel those identified in previous studies of mouse or human cortex (Lake et al., 2016; Tasic et al., 2016), revealing inhibitory neuronal heterogeneity in both cortical layer distribution and the developmental origin from subcortical regions of the medial or caudal ganglionic eminences (MGE or CGE) (Figures 2E–2F and S3C). Therefore, sNucDrop-seq resolves cellular heterogeneity and quantifies cell-type composition at the transcriptomic level with high sensitivity.

sNucDrop-Seq Reveals Layer-Specific Composition and Activity-Dependent Transcriptional State of Cortical Excitatory Neurons

For glutamatergic neuronal clusters, we associated each excitatory neuronal subtype with a combination of known markers indicative of their superficial-to-deep layer distribution (Figure 3A), revealing layer-specific composition of excitatory neuronal subtypes (Figures S3A and S3B). Thus, sNucDrop-seq analysis captures transcriptomic distinctions between closely related subtypes in each cortical layer, which is in high concordance with subtypes previously identified in human (Lake et al., 2016) and mouse (Habib et al., 2017; Tasic et al., 2016; Zeisel et al., 2015) cortices (Figures S3C–S3G).

In response to neuronal activity, excitatory neurons induce expression of hundreds of activity-regulated genes (ARGs), many of which regulate synaptic function and allow neuronal circuits to respond dynamically to experience (Flavell and Greenberg, 2008). Because induced mRNAs may remain in the cytoplasm for hours to days (Schwanhäusser et al., 2011), it is challenging to capture the dynamic and transient neuronal activation process using whole-cell RNA-seq. Direct comparison between single-cell and single-nucleus transcriptomic profiling of activated neurons demonstrated that single-nucleus RNA-seq analysis not only avoids the aberrant activation of ARGs by whole-cell dissociation but also reveals dynamics of transcriptional response to neuronal-activity-inducing experience (Lacar et al., 2016). However, the reported low-throughput single-nucleus RNA-seq analysis requires pre-enrichment of activated neuronal nuclei by sorting and the Fluidigm C1 platform, which is not easily scaled up to accommodate more samples.

On the basis of ARG expression, we explored whether sNucDrop-seq analysis can resolve heterogeneity in activity-induced transcriptional states amongst closely related subtypes. We found that while nuclei in the Ex24 cluster (n=212 nuclei) express nearly identical layer-specific marker genes to E25 (n = 3,628), Ex24 is specifically associated with high-level expression of ARGs (Figures 3A and S4A), including well-defined immediately early genes (IEGs) such as *Fos*, *Arc*, and *Egr1* as well as other activity-regulated transcription factors (e.g., *Npas4*), genes encoding proteins that function at synapses (e.g., *Homer1*), and noncoding RNAs (e.g., *1700016P03Rik*,

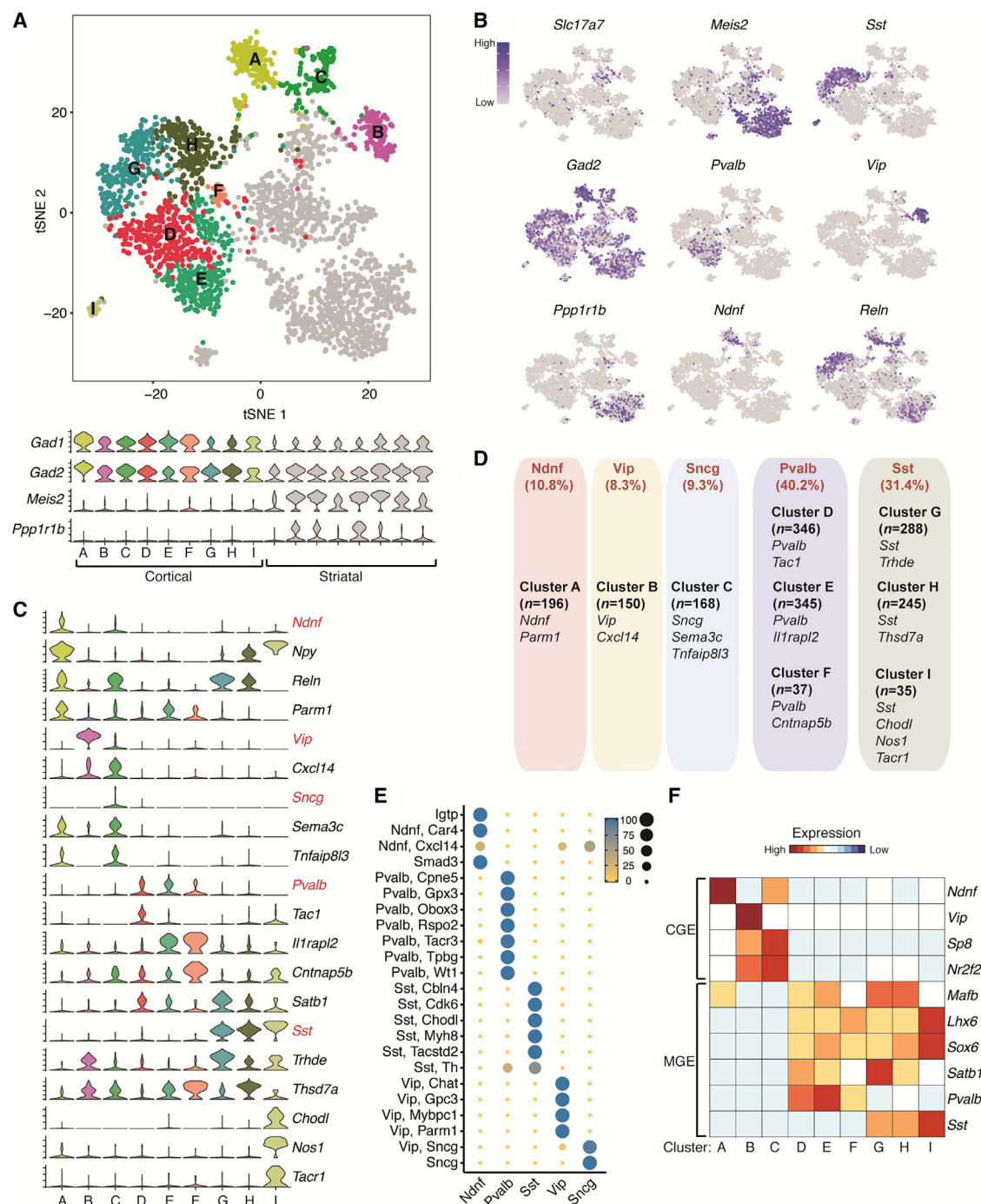


Figure 2. sNucDrop-Seq Reveals Inhibitory Neuronal Subtypes and Composition

(A) Spectral tSNE plot of 1,810 cortical GABAergic neurons and 1,462 noncortical nuclei. The cortical inhibitory neurons are colored (A–H) while noncortical nuclei are marked in grey. Violin plots for two GABAergic neuronal markers (*Gad1* and *Gad2*) and two striatal markers (*Meis2* and *Ppp1r1b*) are shown in the lower panel.

(B) Marker gene expression shown by recoloring and projecting onto the tSNE plot in Figure 2A.

(C) Violin plots showing select marker gene expression for inhibitory neuronal subtypes. Five mutually exclusive subtype-specific marker genes are highlighted in red.

(D) Summary of inhibitory neuronal subtypes identified by sNucDrop-seq. GABAergic subtypes are grouped according to five major classes. Also shown are number of nuclei per subtype and representative marker genes associated with each.

(E) Congruence of five major GABAergic neuronal classes defined by sNucDrop-seq with subpopulations defined from single-cell RNA-seq profiles in the mouse visual cortex (Tasic et al., 2016).

(F) Heatmap showing select marker genes that distinguish cortical inhibitory neurons originated from either CGE or MGE.

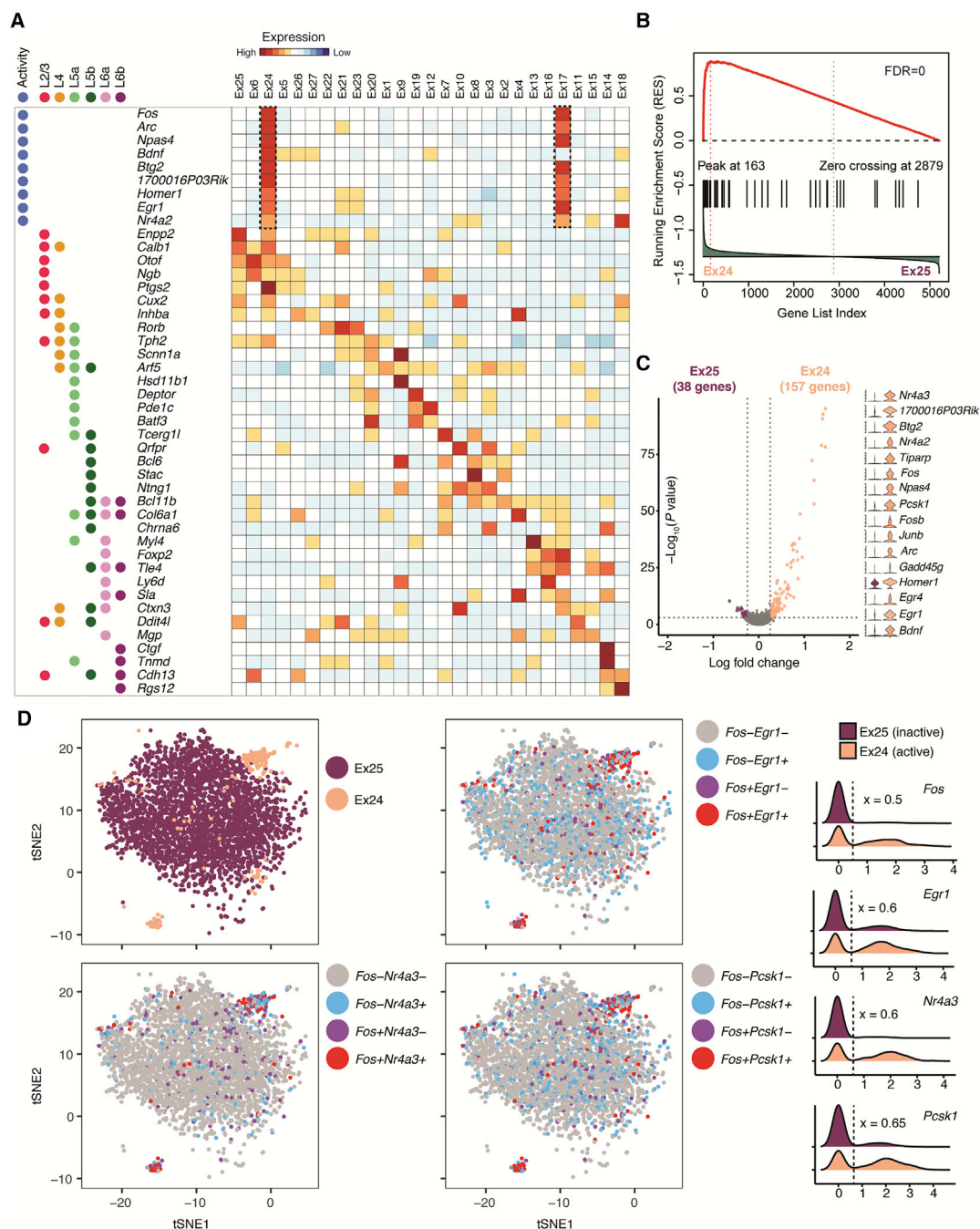


Figure 3. Excitatory Neuronal Subtypes Resolve Heterogeneity in Cortical Layer Distribution and Transcriptional States Driven by Neuronal Activity

(A) Heatmap for layer-specific markers and neuronal-activity-regulated genes showing cortical layer identity (L2/3/, L4, L5a/b, L6a/b), excitatory subtypes, and ARG expression.

(B) GSEA of ARGs between closely related neuronal subclusters. Genes were ranked based on the differential expression between active (left: Ex24) and inactive (right: Ex25). The plot of running enrichment score (RES) is shown in red (top). Vertical bars (in black) in the middle indicate a gene within the ARG gene set. The correlation of gene expression with subclusters is shown in the bottom histogram (in green).

(C) Differential expression between activated and inactivated excitatory neurons within upper (Ex24 versus Ex25)-layer subclusters. Shown on the right are violin plots of select marker gene expression.

(D) Coexpression analysis of neuronal-activity-dependent genes in excitatory neuronal subtypes. Spectral tSNE plots (left) and density plots (right) illustrating expression of select ARGs in upper-layer excitatory neuronal subtypes. The expression level threshold (x) for ARGs was determined *post hoc* and marked by the dash line in density plots. The nuclei in tSNE plots are recolored by ARG expression.

which encodes Mir132). A similar relationship was detected between Ex17 ($n = 91$) and Ex16 ($n = 1,847$). Interestingly, despite variations in number of nuclei and sequencing depth among samples (Table S1), the small percentage of putatively activated neurons (Ex17: $\sim 0.5\%$; Ex24: $\sim 1.2\%$ of all nuclei) was found in nearly all animals (Figure S4B), raising the possibility that neuronal-activity-induced transcriptional states can be reproducibly captured by sNucDrop-seq. We next performed gene set enrichment analysis (GSEA) of a recently curated list of ARGs induced by acute or prolonged neuronal activity (Tyssowski et al., 2017). This analysis indicated that both Ex17 (false-discovery rate [FDR] = 0) and Ex24 (FDR = 0) are significantly enriched for this set of ARGs (Figure 3B). We next determined the genes specifically enriched in activated neurons in Ex17 ($n = 129$ genes as compared to other Ex16) or Ex24 ($n = 157$ genes, as compared to Ex25) neurons (Figure 3C). KEGG pathway analysis suggested that expression signatures of these two excitatory neuronal clusters are enriched for genes involved in the MAPK signaling pathway (adjusted $P = 5.74 \times 10^{-3}$ for Ex17 and 8.14×10^{-3} for Ex24), consistent with previous reports (Lacar et al., 2016; Tyssowski et al., 2017). We also observed heterogeneous expression patterns of ARGs (rapid IEGs: *Fos* and *Egr1*; delayed IEGs: *Nr4a3* and *Pcsk1*) among nuclei in Ex24 (*Fos*⁺: 53% in Ex24 versus 5.7% in Ex25; $P = 1.57\text{E-}71$, Fisher's exact test) and Ex17 (*Fos*⁺: 55% in Ex17 versus 6.8% in Ex17; $P = 3.99\text{E-}31$) excitatory neuronal clusters (Figure 3D and Table S2), in agreement with a continuum of transcriptional states of ARGs revealed by a recent analysis of *Fos*-positive neuronal nuclei isolated from adult mice exposed to a novel environment (Lacar et al., 2016). Together, these results indicate that sNucDrop-seq can potentially identify activity-dependent transcriptional states at single-nucleus resolution *in vivo*.

Mapping Cell-Type-Specific Transcriptional Response to the Pentylentetrazol (PTZ)-Induced Seizure

To further examine cell-type-specific responses to neuronal activity by sNucDrop-seq, we elicited large-scale neuronal activation with PTZ, a GABA(A) receptor antagonist that induces seizures coupled with ARG expression in cortex (Morgan et al., 1987; Yount et al., 1994). Mice were treated with PTZ or received an injection with saline as a control. One hr after injection, nuclei were immediately isolated for sNucDrop-seq analysis (saline: 4,005 nuclei; PTZ: 3,491 nuclei). After filtering out the clusters that did not contain sufficient data points (7 clusters; see STAR Methods), we determined that 15 out of 33 clusters are significantly enriched for the ARG gene set by GSEA (FDR < 0.2) (Figures 4A and 4B). In addition to 11 excitatory neuronal clusters, 2 inhibitory neuronal clusters (Inh2 and Inh5) and 2 non-neuronal clusters (Astro and OPC) were significantly associated with expression of activity-regulated genes in response to PTZ treatment. Our analysis suggests that *Sst*-expressing inhibitory neurons (Inh2) are much more likely to express ARGs than *Pvalb*-expressing neurons (Inh3) in response to PTZ-induced seizure, which is in agreement with a recent cell-type-specific analysis of inhibitory neuronal response to PTZ using a synthetic IEG reporter system (Sørensen et al., 2016). Interestingly, relatively low-abundance interneuron subtypes, such as *Ndnf*-expressing cells (Inh5) and oligodendrocyte precursor cells, also

exhibited significant transcriptional response to PTZ treatment (Figure 4A), suggesting conserved signaling pathways underlying neuronal activation in these cell types.

The activity-dependent gene expression program is likely structured temporally into two major waves of gene activation. The first wave comprises primary response genes (PRGs, also called IEGs). The second wave comprises secondary response genes (SRGs), which require *de novo* translation for their induction and are likely to be regulated by PRG proteins (Tyssowski et al., 2017). Further analysis of top-ranked PTZ-induced ARGs (separated into three groups: rapid PRG, delay PRG, and SRG) in each cluster indicates that rapid PRGs such as *Fos* and *Egr1* are more likely to be detected in inhibitory neuronal clusters (Inh2 and Inh5), whereas delay PRGs (e.g., *Bdnf*, *Mbnl2*) and SRG (e.g., *Nptx2*) are more frequently detected in excitatory neuronal clusters (Figure 4C). This observation suggests that after 1 hr, rapid PRG expression is probably already attenuated in excitatory neurons, and inhibitory neurons are likely activated by the PTZ-induced seizure in a temporally distinct manner. We also found that clusters significantly associated with ARG gene-set induction generally induced a stronger transcriptional response than others (that is, greater number of genes and amplitude of changes) (Figure S4C). Overall, these data should provide a rich resource for identification of genes whose dynamic expression in specific cell types may be important for that cell type's functional response to neuronal activity.

DISCUSSION

In conclusion, sNucDrop-seq is a robust approach for massively parallel analysis of nuclear RNA at single-cell resolution. Because intact nuclear isolation can potentially be accomplished by mechanical douncing and sucrose-gradient ultracentrifugation in almost any primary tissue, including frozen, archived human tissues, sNucDrop-seq and similar approaches (Habib et al., 2017) pave the way to systematically identify cell types, reveal subtype composition, and dissect dynamic functional states such as activity-dependent transcription in complex mammalian tissues.

Our study, with its focus on profiling cortical cells in their native environment and in proportion to their abundance, complements a recent single-cell RNA-seq study of adult mouse visual cortex that deeply surveyed FACS-isolated single cells from a comprehensive collection of transgenic mouse lines (Tasic et al., 2016). We found that the subtypes of both neurons and non-neuronal cells identified in the two studies are highly consistent (Figure S3C). Overall, our unbiased sampling approach may provide a more complete and accurate description of the cell-type landscapes, whereas deep sequencing of full-length mRNAs from select cells may identify a more complete set of molecular markers for each subtype.

In addition to neuronal subtype classification, precise localization and quantification of the activity of all neurons across specific brain regions would provide insights into how neural information is processed in physiological and pathological states. In mammals, a snapshot of the activity of ensembles of neurons in the mammalian brain can be obtained by whole-brain tissue

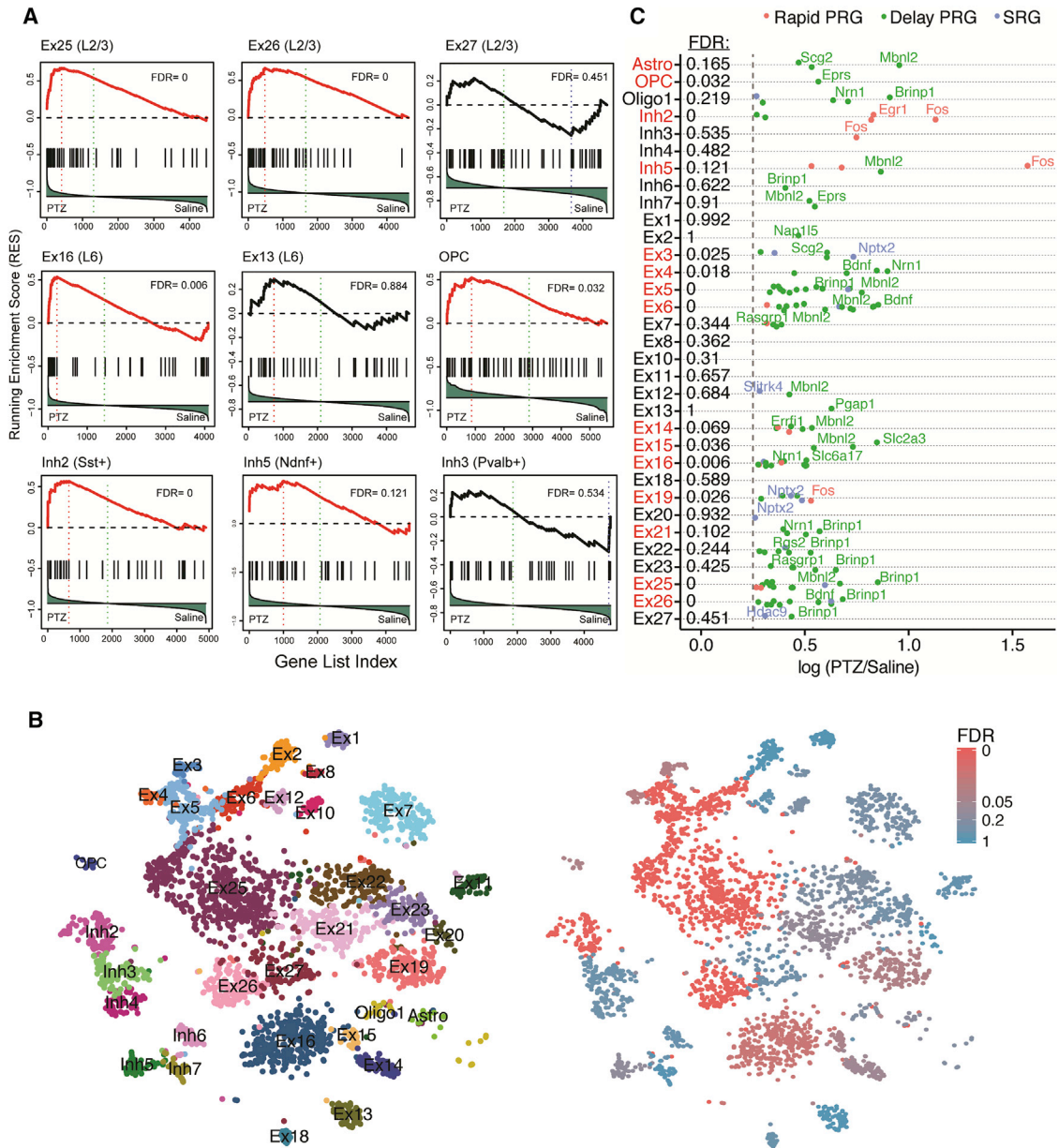


Figure 4. sNucDrop-Seq Reveals Cell-Type Specific Transcriptional Responses to PTZ-Induced Seizure

(A) GSEA of the ARG set in different cell clusters by the comparison between PTZ- and saline-treated neurons was performed as in [Figure 3B](#). For each cell type, genes were ranked based on the differential expression between PTZ (left) and saline (right). The FDR for GSEA of ARGs is listed, and clusters with FDR < 0.2 are highlighted in red.

(B) Spectral tSNE plot of 3,491 nuclei from PTZ-treated mouse cortex. The nuclei were colored per annotated cell types (left) or the FDR of GSEA analysis (right). (C) Cell-type-specific induction of ARGs in response to PTZ treatment. Rapid PRGs, delayed PRGs, and SRGs are highlighted in red, green, and blue, respectively. The x axis indicates the natural log fold change of gene expression between PTZ and saline. The dashed line denotes the cutoff of gene expression fold change (log 0.25). The FDR for GSEA of ARGs (as in Figure 4A) is listed, and clusters with FDR < 0.2 are highlighted in red.

clearing coupled with immunolabeling of IEGs (Renier et al., 2016), which requires advanced instrumentation and has low temporal resolution (that is, IEG transcripts can outlast the end of activity by over 4 hr). Single-nucleus RNA-seq approaches such as sNucDrop-seq rely on isolation of nuclei, thereby enriching nascent transcripts and enabling the detection and quantification of neural activity (using expression of ARGs as proxies) at

much higher temporal resolution. In addition, our results show that transcriptomic differences between closely related subtypes (e.g., between Ex24 and Ex25) may be largely driven by neuronal-activity-dependent transcriptional programs rather than by differences in developmental origin or anatomical distribution. This observation suggests that environmental stimuli may contribute to discontinuous transcriptomic differences.

and future studies are needed to investigate how continuous variables, such as neuronal activity, can be best implemented as additional classifiers for cell types and functional states.

LIMITATIONS

As with other single-cell or single-nucleus RNA-seq approaches, sNucDrop-seq relies on dissociation of individual cells or nuclei in dissected tissues, thereby discarding cells' original anatomical location information. Thus, combining a sequencing-based single-cell genomics approach with imaging-based multiplex fluorescence *in situ* hybridization (FISH) methods (Chen et al., 2016; Chen et al., 2015; Shah et al., 2016) that allow detection of a set of preselected genes in tissues is a promising approach to acquire essential information about precise anatomical location in order to reconstruct a more complete cell-type atlas in complex mammalian tissues.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental Information includes four figures, two tables, three videos, and supplemental methods and can be found with this article online at <https://doi.org/10.1016/j.molcel.2017.11.017>.

AUTHOR CONTRIBUTIONS

H.W. and Z.Z. conceived the study. H.W., P.H., E.F., D.K., and S.T. performed experiments. H.W. and P.H. carried out data analysis. H.W. wrote the manuscript with input from other authors.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, Peptides, and Recombinant Proteins		
Dulbecco's Modified Eagle's Medium	Life Technologies	11965084
Fetal Bovine Serum	Life Technologies	26140079
L-glutamine	Life Technologies	25030081
0.05% Trypsin	Life Technologies	25300054
Matrigel matrix	Corning	354230
DMEM/F12	Life Technologies	11320033
TeSR-E8 Medium	Stem Cell Technologies	05940
DPBS, no calcium, no magnesium	Invitrogen	14190136
Sucrose	Sigma-Aldrich	S0389-1KG
1M Tris-HCl, pH 8.0	Invitrogen	15568-025
MgAc ₂	Sigma-Aldrich	M5661-50G
cOmplete™, EDTA-free Protease Inhibitor Cocktail	Roche	11873580001
CaCl ₂	Sigma-Aldrich	C1016-500G
Triton X-100	Sigma-Aldrich	T8787-100mL
0.5M EDTA, pH 8.0	Invitrogen	15575-020
NxGen RNase Inhibitor	Lucigen	30281-2
Bovine Serum Albumin	Sigma-Aldrich	A8806-5G
Ficoll PM-400	GE Healthcare/Fisher Scientific	45-001-745
Sarkosyl	Sigma-Aldrich	L7414-50mL
DTT	Fermentas	R0862
QX200 Droplet Generation Oil for EvaGreen	Bio-Rad	186-4006
Perfluoro-1-octanol	Sigma-Aldrich	370533-25G
dNTPs	Clontech	639125
Critical Commercial Assays		
Maxima H Minus Reverse Transcriptase	ThermoFisher	EP0753
KAPA HiFi hotstart readymix	KAPA Biosystems	KK2602
Deposited Data		
Raw and analyzed data	This paper	GEO: GSE106678
Experimental Models: Cell Lines		
NIH3T3	ATCC	CRL-1658
H7 (female, human embryonic stem cells)	WiCell	WA07
Experimental Models: Organisms/Strains		
Mouse: C57BL/6J	Jackson Laboratory	000664
Oligonucleotides		
Template Switch Oligo: AAGCAGTGGTATCAACGC AGAGTGAATrGrGrG	Macosko et al., 2015	N/A
TSO-PCR primer: AAGCAGTGGTATCAACGCAGAGT	Macosko et al., 2015	N/A
P5-TSO hybrid primer: AATGATACGGCGACCAACCG AGATCTACAGCCTGTCCGCGGAAGCAGTGGTAT CAACGCAGAGT*A*C	Macosko et al., 2015	N/A
Custom Read 1 Primer: GCCTGTCCGCGGAAGCA GTGGTATCAACGCAGAGTAC	Macosko et al., 2015	N/A

(Continued on next page)

Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
Drop-seq_tools (v1.12)	http://mccarrolllab.com/dropseq/	N/A
STAR v2.5.2a	https://github.com/alexdobin/STAR	N/A
Seurat v1.4	http://satijalab.org/seurat/	N/A
Seurat v2.0	http://satijalab.org/seurat/	N/A
GSEA	http://software.broadinstitute.org/gsea/index.jsp	N/A
DBSCAN	https://cran.r-project.org/web/packages/dbscan/index.html	N/A
MISO	https://miso.readthedocs.io/en/fastmiso/	N/A
Random Forest	https://www.stat.berkeley.edu/~breiman/RandomForests/	N/A
Other		
Detailed Bench Protocol	This paper	Methods S1
Tube, Thinwall, Polypropylene, 38.5 mL, 25 x 89 mm (qty. 50)	Beckman Coulter	326823
Glass 15mL Dounce Tissue Grinder Set with Two Glass Pestles, Grinding Chamber O.D. x L: 22 x 94mm (Case of 2)	Wheaton	357544
SW 28 Ti Rotor, Swinging Bucket, Aluminum, 6 x 38.5 mL, 28,000 rpm, 141,000 x g	Beckman Coulter	342207
Barcoded Beads	ChemGenes	MACOSKO-2011-10
PDMS Microfluidic Device	μFluidix	Batch #9508
Syringe Pumps	KD Scientific	78-8100
40μm Sterile Cell Strainer	Fisher Scientific	22-363-547
Medical Grade Polyethylene Micro Tubing	Scientific Commodities	BB31695-PE/2
SPRISelect Beads	Beckman Coulter	B23318
75-cycle High Output v2 Kit	Illumina	FC-404-2005
10-micron carboxylated polystyrene beads	Bangs Labs	#PC06N-11355

CONTACT FOR REAGENT AND RESOURCE SHARING

Requests should be addressed to and will be fulfilled by Lead Contact Hao Wu (haowu2@pennmedicine.upenn.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS**Cell Lines and Culture Conditions**

Mouse NIH3T3 cells were purchased from ATCC and were grown in Dulbecco's Modified Eagle's Medium (DMEM) (Life Technologies) supplemented with 10% fetal bovine serum (FBS) (Life Technologies) and 2 mM L-glutamine (Life Technologies) at 37°C in 5% CO₂. The culture was passaged every 2-3 days using 0.05% Trypsin (Life Technologies). Female, human embryonic stem cells (H7, NIH registration number 0061) have been procured from WiCell (Madison, WI) and maintained at 37°C in 5% CO₂ on growth factor-reduced Matrigel matrix (Corning) coated six-well tissue culture plates. The six-well plates were coated with diluted (1:30) Matrigel matrix in DMEM/F12 (Life Technologies). H7 cells (between passage 40 and 70) were maintained in TeSR-E8 medium (Stem Cell Technologies) and passaged every 5-6 days as small aggregates using an enzymatic digestion-free method (0.5 mM EDTA in DPBS without CaCl₂ and MgCl₂ [Life Technologies]).

Animals

Experiments were conducted in accordance with the ethical guidelines of the National Institutes of Health and with the approval of the Institutional Animal Care and Use Committee of the University of Pennsylvania. Mice were group-housed in cages of three to five in a 12-hr light/dark cycle with food and water provided ad libitum. All mice used for experiments were naive to behavioral assays and other procedures. For pentylenetetrazole (PTZ)-induced seizures, 10-week old, wild type, male F1 mice from a C56BL/6J to FVB/NJ

breeding were used. All others were 6- to 10-week-old, wild type, male mice in a pure C57BL/6J background. For PTZ treatment, mice were injected intraperitoneally with PTZ at 50 mg/kg body weight or received an injection with saline as a control. One hr after injection, the mice were sacrificed.

METHOD DETAILS

Isolation and Purification of Nuclei

Mouse brains (postnatal 6-10 weeks) were rapidly resected on ice. Cortices were freshly processed or flash frozen in liquid nitrogen for 2 minutes and subsequently kept at -80°C before nuclear isolation. Nuclei were isolated and purified as previously described with some modifications (Johnson et al., 2017). Briefly, 14 mL of sucrose cushion (1.8 M sucrose (Sigma-Aldrich), RNase & DNase free, ultra pure grade), 10 mM Tris-HCl pH 8.0 (Invitrogen), 3 mM MgAc_2 (Sigma-Aldrich), protease inhibitor cocktail (Roche) was added to the bottom of centrifuge tubes (Beckman Coulter). Using a glass homogenizer (Wheaton), a freshly isolated or frozen mouse cortex sample was subjected to dounce homogenization (21 times with loose pestle followed by 7 times with tight pestle) in 12 mL of homogenization buffer (0.32M sucrose, 5 mM CaCl_2 (Sigma-Aldrich), 3mM MgAc_2 , 10 mM Tris-HCl pH 8.0, 0.1% Triton X-100 (Sigma-Aldrich), 0.1 mM EDTA (Invitrogen), protease inhibitor cocktail). For *in vitro* cultured cells, cell pellets (~ 5 million cells) were resuspended in homogenization buffer and dounced 20 times with a loose pestle. Homogenates (~ 12 mL) were layered onto the sucrose cushion in the centrifuge tubes, and 10 mL of homogenization buffer was added atop of the homogenates. The tubes were then centrifuged in a Beckman Coulter L7-65 Ultracentrifuge at 25,000 rpm at 4°C for 2 hr using a Beckman Coulter SW28 swinging bucket rotor (Beckman Coulter). The supernatant was carefully removed via aspiration. See [Methods Video S1](#) for details of how nuclei are harvested upon sucrose-gradient ultracentrifugation. 1 mL of chilled DPBS with RNase inhibitor (Lucigen) was added to resuspend the nuclear pellet, and nuclei were subsequently transferred to a 1.5-mL tube. Nuclei were pelleted at 5,000 rpm for 10 min at 4°C , and then resuspended in 0.01% BSA (Sigma-Aldrich) in DPBS. See [Methods Video S2](#) for details of nuclei pelleting, resuspension to form single-nucleus suspension, and nuclear counting. After resuspension, nuclei were filtered through a 40- μm cell strainer (Fisher Scientific), visually inspected for morphology and quality assurance, and counted using a Fuchs-Rosenthal counting chamber before droplet microfluidic encapsulation. For mouse cortices, we obtain $3.45 \pm 2.00 \times 10^6$ nuclei, per round of isolation (based on 11 measurements). The nuclear isolation efficiency for *in vitro* cultured cells is $\sim 84\%$ (number of nuclei/number of input cells $\times 100$).

Single-Nucleus RNA-Seq Library Preparation and Sequencing

The nuclear suspension was diluted to a concentration of 100 nuclei/ μL in DPBS containing 0.01% BSA. Approximately 1.25 mL of this single-nucleus suspension was loaded for each sNucDrop-seq run. The single-nucleus suspension was then co-encapsulated with barcoded beads (ChemGenes) using an Aquapel-coated PDMS microfluidic device ($\mu\text{Fluidix}$) connected to syringe pumps (KD Scientific) via polyethylene tubing with an inner diameter of 0.38 mm (Scientific Commodities). See [Methods Video S3](#) for an overview of the microfluidics setup used to co-encapsulate single nuclei with barcoded beads. Barcoded beads were resuspended in lysis buffer (200 mM Tris-HCl pH8.0, 20 mM EDTA, 6% Ficoll PM-400 (GE Healthcare/Fisher Scientific), 0.2% Sarkosyl (Sigma-Aldrich), and 50 mM DTT (Fermentas; freshly made on the day of run) at a concentration of 120 beads/ μL . The flow rates for nuclei and beads were set to 4,000 $\mu\text{L/hr}$, while QX200 droplet generation oil (Bio-rad) was run at 15,000 $\mu\text{L/hr}$. A typical run lasts ~ 20 min. Droplet breakage with Perfluoro-1-octanol (Sigma-Aldrich), reverse transcription and exonuclease I treatment were performed, as previously described, with minor modifications (Macosko et al., 2015). For up to 120,000 beads, 200 μL of reverse transcription (RT) mix (1x Maxima RT buffer (ThermoFisher), 4% Ficoll PM-400, 1 mM dNTPs (Clontech), 1 U/ μL RNase inhibitor, 2.5 μM Template Switch Oligo (AAGCAGTGGTATCAACGCAGAGTGAATrGrGrG) (Macosko et al., 2015), and 10 U/ μL Maxima H Minus Reverse Transcriptase (ThermoFisher)) were added. The RT reaction was incubated at room temperature for 30 minutes, followed by incubation at 42°C for 150 minutes. To determine an optimal number of PCR cycles for amplification of cDNA, an aliquot of 6,000 beads (corresponding to ~ 100 nuclei for a capture rate of 1.5–2%) was amplified by PCR in a volume of 50 μL (25 μL of 2x KAPA HiFi hotstart readymix (KAPA biosystems), 0.4 μL of 100 μM TSO-PCR primer (AAGCAGTGGTATCAACGCAGAGT (Macosko et al., 2015), 24.6 μL of nuclease-free water) with the following thermal cycling parameter (95°C for 3 min; 4 cycles of 98°C for 20 sec, 65°C for 45 sec, 72°C for 3 min; 9 cycles of 98°C for 20 sec, 67°C for 45 sec, 72°C for 3 min; 72°C for 5 min, hold at 4°C). After two rounds of purification with 0.6x SPRISelect beads (Beckman Coulter), amplified cDNA was eluted with 10 μL of water. 10% of amplified cDNA was used to perform real-time PCR analysis (1 μL of purified cDNA, 0.2 μL of 25 μM TSO-PCR primer, 5 μL of 2x KAPA FAST qPCR readymix, and 3.8 μL of water) to determine the additional number of PCR cycles needed for optimal cDNA amplification (Applied Biosystems QuantStudio 7 Flex). We then prepared PCR reactions per total number of barcoded beads collected for each sNucDrop-Seq-seq run, using 6,000 beads per 50- μL PCR reaction, and ran the aforementioned program to amplify the cDNA for 4 + 10 to 12 cycles. We then tagged cDNA using the Nextera XT DNA sample preparation kit (Illumina, cat# FC-131-1096), starting with 550 pg of cDNA pooled in equal amounts, from all PCR reactions for a given run. Following cDNA tagmentation, we further amplified the tagmented cDNA libraries with 12 enrichment PCR cycles using the Illumina Nextera XT i7 primers along with the P5-TSO hybrid primer (AATGATACGGCGACCACCGAGATCTACACGCTGTCCGCGGAAGCAGTGGTATCAACGCAGAGT*A*C) (Macosko et al., 2015). After quality control analysis by Qubit 3.0 (Invitrogen) and a Bioanalyzer (Agilent), libraries were sequenced on an Illumina

NextSeq 500 instrument using the 75-cycle High Output v2 Kit (Illumina). We loaded the library at 2.0 pM and provided Custom Read1 Primer (GCCTGTCCGCGGAAGCAGTGGTATCAACGCAGAGTAC) at 0.3 μ M in position 7 of the reagent cartridge. The sequencing configuration was 20 bp (Read1), 8 bp (Index1), and 50 or 60 bp (Read2). In total, 17 mouse cortex samples were analyzed with sNucDrop-seq in four sequencing runs.

Single-Cell RNA-Seq Library Preparation and Sequencing

Drop-seq was performed as previously described, with default settings (Macosko et al., 2015). The cell suspension was diluted to 100 cells/ μ L with DPBS containing 0.01% BSA and 1 mL cell suspension was loaded for each Drop-seq run. After cell capture, reverse transcription, exonuclease treatment, cDNA amplification and tagmentation, libraries were diluted to 1.9 pmol, and equal amounts of multiplexed libraries were mixed and subjected to paired-end sequencing on Illumina NextSeq 500 sequencer as described for sNucDrop-seq. 10x Genomics single-cell 3' libraries were constructed, as previously described (Zheng et al., 2017), with recommended settings using Chromium single cell 3' v2 reagent kits by the Center for Applied Genomics of The Children's Hospital of Philadelphia (CHOP). It is worth noting that the 10x Genomics single-cell 3' solution workflow supports cDNA amplification only from whole cells, possibly due to inefficient lysis of nuclei purified via sucrose-gradient ultracentrifugation. 10x genomics recently released a specific nucleus isolation and purification protocol (<https://support.10xgenomics.com/single-cell-gene-expression/index/>) that may support single-nucleus RNA-seq analysis on the 10x genomics platform.

Microfluidics Device for sNucDrop-Seq

To examine that the channel depth of the microfluidics device (custom-made by μ Fluidix) used for sNucDrop-seq, droplet volume was measured using 10 μ m carboxylated polystyrene fluorescent beads (Bangs Labs), as previously described (<http://mccarrolllab.com/dropseq/>). The beads were first suspended at a concentration of 1,000 beads/ μ L in lysis buffer (200 mM Tris-HCl pH8.0, 20 mM EDTA, 6% Ficoll PM-400, 0.2% Sarkosyl, and 50 mM DTT). Drop-seq was performed as previously described, with default settings (Macosko et al., 2015), except the syringe pump intended for cells was loaded with DPBS. The total number of beads encapsulated in several hundred droplets was determined, and droplet volume was calculated, as follows:

$$\text{Droplet volume} = \left(\frac{\text{Total number of beads, inside droplets, counted}}{\text{Total number of droplets counted}} \right) / 500 \text{ beads per microliter}$$

We then determined the droplet volume (1.2 nL) and estimated the radius of droplets (1/2 of channel depth) as 65 μ m. This confirms that the channel depth of microfluidics devices used for sNucDrop-seq experiments is approximately \sim 130 μ m.

It is worth noting that while both sNucDrop-seq (this study) and DroNc-seq (Habib et al., 2017) have adapted the Drop-seq workflow for single-nucleus RNA-seq analysis of mammalian brains, there are several technical differences between the two. First, different nuclei purification methods were employed. sNucDrop-seq utilizes home-brew suffers for sucrose-gradient ultracentrifugation, whereas DroNc-seq relies on a commercially available kit. Second, while sNucDrop-seq utilizes standard Drop-seq microfluidics devices (channel depth: 125 μ m), DroNc-seq requires a modified design (channel depth: 75 μ m). Because of the reduced channel depth, the DroNc-seq specific device is more likely to clog and requires pre-filtering of bar-coded beads to enrich beads of smaller sizes (causing 50% loss of this expensive reagent). Thus, while sNucDrop-seq requires special instrument such as ultracentrifuge, we estimate that sNucDrop-seq is more cost-effective than either DroNc-seq or 10x Genomics platform.

QUANTIFICATION AND STATISTICAL ANALYSIS

Preprocessing of sNucDrop-Seq Data

Paired-end sequencing reads of sNucDrop-seq were processed using publicly available the Drop-seq Tools v1.12 software (Macosko et al., 2015) with some modifications. Briefly, each mRNA read (read2) was tagged with the cell barcode (bases 1 to 12 of read 1) and unique molecular identifier (UMI, bases 13 to 20 of read 1), trimmed of sequencing adaptors and poly-A sequences, and aligned using STAR v 2.5.2a to the mouse (mm10, Gencode release vM13) or a concatenation of the mouse and human (for the species-mixing experiment) reference genome assembly. Because a substantial proportion (\sim 50%) of reads derived from nuclear transcriptomes of mouse cortices were mapped to the intronic regions, the intronic reads were retained for downstream analysis. A custom Perl script was used to retrieve both exonic and intronic reads mapped to predicted strands of annotated genes. Uniquely mapped reads were grouped by cell barcodes. Cell barcodes were corrected for possible bead synthesis errors, using the *DetectBeadSynthesisErrors* program from the Drop-seq Tools v1.12 software. To generate digital expression matrix, a list of UMIs in each gene (as rows), within each cell (as columns), was assembled, and UMIs within ED = 1 were merged together. The total number of unique UMI sequences was counted, and this number was reported as the number of transcripts of that gene for a given nucleus.

Comparison of Drop-Seq (Whole Cells) and sNucDrop-Seq (Nuclei)

Only mouse NIH3T3 cells or nuclei that expressed >800 genes were retained for analysis. For each gene, the average normalized expression level was calculated as $\log(\text{normalized UMI counts} + 1)$.

Clustering and Marker Gene Identification

Raw digital expression matrices were combined and loaded into the R package Seurat. For normalization, UMI counts for all nuclei were scaled by library size (total UMI counts), multiplied by 10,000 and transformed to log scale. Only genes found to be expressing in >10 cells were retained. Nuclei with a relatively high percentage of UMIs mapped to mitochondrial genes (≥ 0.1) were discarded. Moreover, nuclei with fewer than 800 or more than 6,000 detected genes were discarded, resulting in 20,858 nuclei that pass filter.

Before clustering, batch effects from multiple animals were regressed out using the function *RegressOut* in R package Seurat v1.4 (Macosko et al., 2015). The highly variable genes were identified using the function *MeanVarPlot* with the parameters: *x.low.cutoff* = 0.0125, *x.high.cutoff* = 3 and *y.cutoff* = 0.8, resulting in an output of 1,932 highly variable genes. The expression level of highly variable genes in the nuclei was scaled and centered along each gene, and was conducted to principal component analysis. We then used two methods to assess the number of PCs to be utilized in downstream analysis: 1) The cumulative standard deviations of each PC were plotted using the function *PCElbowPlot* in Seurat to identify the ‘knee’ point at a PC number after which successive PCs explain diminishing degrees of variance, and 2) the significance for each gene’s association with each PC was accessed by the function *JackStraw* in Seurat. Based on these two methods, we selected first 30 PCs for two-dimensional t-distributed stochastic neighbor embedding (tSNE), implemented by the Seurat software with the default parameters. Based on the tSNE map, twenty-one clusters were identified using the function *FindCluster* in Seurat with the resolution parameter set to 2. Clusters that co-express both non-neuronal and neuronal markers, representing cell doublets, were removed. Based on expression of well-established marker genes, we assigned 19,241 nuclei to cerebral cortex (93.1% of our data) and 1,415 nuclei to non-cortical neurons (6.9%). To perform clustering analysis on 19,241 cortical cells, we identified 1,968 highly variable genes and selected first 40 PCs. The nuclei were classified into 44 clusters with the resolution parameter set to 4. To identify clusters with highly similar identity, we performed pairwise comparison to identify differentially expressed genes using the function *FindMarkers* in Seurat, with likelihood-ratio test. We merged the clusters exhibiting less than 10 genes with an average expression difference greater than 2-fold between clusters. Based on two-dimensional coordinates of nuclei in the tSNE plot, a density-based clustering method implemented in the DBSCAN R package was used to identify outliers, filtering out 882 nuclei (4.5% of 19,241 nuclei) with the reachability distance parameter (*eps*) setting to 0.9 and minimal number of nuclei within that *eps* radius setting to 10. Finally, we excluded clusters containing expression markers for more than one canonical cell type. As a result of these steps, we were able to assign 18,194 (94.6% of cortical nuclei) nuclei into 40 clusters.

To identify the marker genes, differential expression analysis was performed by the function *FindAllMarkers* in Seurat with likelihood-ratio test. Differentially expressed genes that were expressed at least in 25% cells within the cluster and with a fold change more than 0.5 (log scale) were considered to be marker genes. In total, 2,001 protein-coding genes and 90 long noncoding RNAs were identified for 40 clusters. For the marker genes, average gene expression for each cluster was determined, and Euclidean distances between all pairs was calculated. This dataset was used as input for complete linkage hierarchical clustering and dendrogram assembly. To generate a heatmap of marker genes across clusters, the average expression level of marker genes within each cluster were calculated. For each cluster, the average expression was centered and scaled by each gene. Next, the *hclust* function in the R was used to generate the cluster dendrogram with the “ward.D” method.

Sub-clustering of Cortical and Non-cortical Inhibitory Neurons

Cortical inhibitory neuronal nuclei from cluster *Inh* 1–7 were first combined with non-cortical nuclei. We then identified 2,139 highly variable genes in a total of 3,425 nuclei and performed PCA analysis. The significance of PCs was examined by the *Jackstraw* function in Seurat. First 23 significant PCs with *P*-value < $1\text{E-}8$ were selected for clustering analysis and 19 sub-clusters were identified with the resolution setting to 2. After merging clusters, identifying/removing outliers (DBSCAN), and filtering cell doublets, we identified 9 cortical inhibitory neuronal sub-groups (1,810 nuclei), and 8 sub-clusters for non-cortical nuclei (1,462 nuclei).

Identification of Differentially Expressed Genes

Differential gene expression analysis between PTZ treatment and saline groups or between active and inactive neurons was performed using the function *FindMarkers* in Seurat, using a likelihood ratio test. For the PTZ and saline comparison, genes with a fold-change of more than 0.25 (log scale) and a *P*-value less than 0.01 were considered to be differentially expressed, while genes showing 0.25-fold change (log-scale) with *P*-value < $1\text{E-}3$ were defined as differentially expressed between active and inactive neurons. For the PTZ and saline comparison within each cluster, only the clusters containing at least 20 nuclei in both two conditions, PTZ and saline, were tested.

Gene Set Enrichment Analysis of Neuronal-Activity-Regulated Genes (ARGs)

The list of 172 ARGs was previously defined (Tyssowski et al., 2017). These ARGs were classified into 3 groups: rapid primary response genes (rapid PRG), delayed primary response genes (delayed PRG) and secondary response genes (SRG) based on the expression pattern across the time points in response to the KCl stimulation of cultured primary neurons (Tyssowski et al., 2017). For each cluster, only genes that were expressed in at least of 10% nuclei were considered and the expression matrix of those genes

in each cluster was loaded into the GSEA analysis (<http://software.broadinstitute.org/gsea/index.jsp>). Clusters with an FDR < 0.2 were considered significant.

Alternative Splicing Analysis

To identify the cell-type-specific alternative splicing events, we classify the 40 clusters into 8 cell types: excitatory neuronal layer 2/3 (Cluster Ex5, Ex6, Ex24, Ex25, Ex26, and Ex27), excitatory neuronal layer 4 (cluster Ex21, Ex22, and Ex23), excitatory neuronal layer 5/6 (Cluster Ex1, Ex2, Ex3, Ex4, Ex7, Ex8, Ex9, Ex10, Ex12, Ex19 and Ex20), excitatory neuronal layer 6 (Cluster Ex11, Ex13, Ex14, Ex15, Ex16, Ex17 and Ex18), inhibitory neuronal Sst+ (Cluster Inh1 and Inh2), inhibitory neuronal Pvalb+ (Cluster Inh3 and Inh4), other inhibitory neurons (Cluster Inh5, Inh6 and Inh7), and non-neuronal cell types (Cluster EC, MG, Astro, OPC, Oligo1 and Oligo2). The bam files of the nuclei from the same cell type were merged and sorted by Samtools. Pairwise comparison of differential splicing events between cell types was performed using MISO (Katz et al., 2010). Splicing events with a bayes factor (>10), ΔPSI >0.2; at least 1 read supports the inclusion or exclusion isoform, and at least 10 reads supports either of these events.

Comparison of sNucDrop-Seq Data to Other Single-Cell/Nucleus RNA-Seq Data Sets

To assess the validity of sNucDrop-seq results, we compare our data with recently published single-nucleus RNA-seq data from mouse prefrontal cortex (Habib et al., 2017) or single-cell RNA-seq data derived from mouse visual cortex (Tasic et al., 2016). To compare cell-type-specific expression signatures defined by sNucDrop-seq and those of other data sets, we computed the pairwise Pearson correlation coefficients between each pair of cell types in other data sets and our sNucDrop-seq data set for a common set of genes. To generate a common marker gene list for the pairwise comparison, we identified 1,986 (95.0%) and 1,919 (91.8%) common marker genes for the DroNuc-seq data set (Figure 1G) and the single-cell data set (Figure S3C), respectively. Average natural log scaled UMI counts or TPM counts were used to generate the DroNuc-seq (Habib et al., 2017) and single-cell RNA-seq (Tasic et al., 2016) gene expression matrix, respectively. R function *cor.test* was used to calculate the pairwise Pearson correlation coefficients.

To further examine whether GABAergic neuronal subclusters identified in this study agree with a previous single-cell RNA-seq study of well-characterized transgenic mouse lines (Tasic et al., 2016), we adopted a random forest classifier-based method (Shekhar et al., 2016) (Figure 2E). First, we used five major GABAergic neuron sub-types defined by sNucDrop-seq to train a random forest classifier on our sNucDrop-seq dataset. We first sampled 60% of nuclei from each cell type to build the training set of 1,204 nuclei, and the remaining 40% nuclei (770) were used as test set for evaluating the performance of the trained classifier. We then identified the 1,182 most highly variable genes using the Seurat function *MeanVarPlot* with the following parameters: *x.low.cutoff* = 0.0125, *x.high.cutoff* = 4 and *y.cutoff* = 0.8. The digital expression matrix comprising 1,182 genes across 1,204 nuclei was used to build the classifier. We trained a random forest classifier using the R package *randomForest* with the *ntree* parameter set to 1,000. By applying this classifier to assign the nuclei in the test set, 96.9% of the nuclei were correctly mapped to their classes. Then the classifier was used to map the cells in single cell dataset from transgenic Cre lines (Tasic et al., 2016).

By utilizing a recently published computational strategy for integrated analysis of multiple data sets (implemented in Seurat v2.0), we further performed comparative analysis of the DroNuc-seq data set (Habib et al., 2017) and this study (Figures S3E–S3G). First, we downloaded the raw DroNuc-seq data (mouse prefrontal cortex) and generated the digital expression matrix using our pipeline that utilizes both exonic and intronic reads. After filtering low quality nuclei, 5,441 nuclei from DroNuc-seq and all 18,194 nuclei from our dataset with > 800 genes expressed were imported into Seurat. Then, the union of the top 1,000 genes with the highest dispersion from both datasets was subject to a canonical correlation analysis (CCA) to identify common sources of variation between the two datasets. The first 15 dimensions of the CCA was chosen to align the two datasets. Nuclei whose expression level cannot be explained by those 15 CCA were removed. The remaining nuclei, that passed filter, were subject to clustering with the resolution parameter set to 1.2. We assigned 23,213 (98.2% of input data) comprised 5,337 DroNuc-seq nuclei and 17,876 scNucDrop-seq nuclei into 18 clusters.

Identification of Nuclei Expressing ARGs

To evaluate the fraction and heterogeneity of ARG-expressing nuclei in cortical excitatory neurons (Figure 3D), we examined the co-expression pattern of 4 selected ARGs (rapid PRGs: *Fos* and *Egr1*; delayed PRGs: *Nr4a3* and *Pcsk1*). Based on the distribution of gene expression levels between active (Ex24) and inactive (Ex25) neurons, we determined the baseline expression level of these ARGs, *post-hoc*. The nuclei whose expression level was higher than the baseline ARG expression level were classified as ARG-expressing nuclei. We then examined whether the proportions of ARG-expressing nuclei were higher in the active neurons compared to the inactive neurons using Fisher's exact test (Table S2).

To compare the proportion of ARG-expressing nuclei across all the cell types (Figure S4A), mean and standard deviation (SD) of selected ARGs were computed. Nuclei whose expression level of ARGs was higher than mean+2SD are defined as ARG-expressing nuclei. Then the percentage of ARG-expressing nuclei was reported for each cluster, defined in Figure 1C.

DATA AND SOFTWARE AVAILABILITY

The next-generation sequencing data reported in this study have been deposited to the Gene Expression Omnibus (GEO). The accession number for this data is GEO: GSE106678. R scripts for data analysis are available on GitHub repository (https://github.com/wulabupenn/Hu_MolCell_2017).

ADDITIONAL RESOURCES

Detailed Protocol

A detailed bench protocol is supplied in an attached file, entitled [Methods S1](#).